

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Letters Patent of:
Michael HALLEK et al.

Patent No.: 7,314,912

Issued: January 1, 2008

For: AAV STRUCTURAL PROTEIN, ITS
PREPARATION AND USE

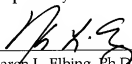
REQUEST FOR CERTIFICATE OF CORRECTION
PURSUANT TO 37 C.F.R. § 1.322

Upon reviewing the above-identified patent, Patentee noted typographical and clerical errors made by the US Patent Office which should be corrected. The errors to be corrected are described in detail on the enclosed PTO Form 1050.

Applicant believes no fee is due with this request, as the errors to be corrected were made by the PTO. However, if a fee is due, please charge our Deposit Account No. 03-2095, under Order No. 50796-016001 from which the undersigned is authorized to draw.

Respectfully submitted,

Date: 22 August 2012



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UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. 7,314,912
DATED Jan. 1, 2008
INVENTORS Hallek et al.

It is certified that an error appears or errors appear in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

On Cover Page, under U.S. PATENT DOCUMENTS, replace
"2001/0031463" with --2001/031463--.

Column 2, Line 21, replace "determined are: are:" with
--determined are:--;

Line 37, replace "AVV" capsid" with --AAV capsid--.

Column 5, Line 36, replace "AAV" with --AAV structural protein
which is located, for example, on a helper plasmid. Packaging
with the mutant helper plasmid results in recombinant AAV with P1
in the capsid (rAAV-P1).--

Column 8, Line 61, replace "YYLSR" with --YYLSR--;

Line 62, replace "EEKFF" with --EEKFF--;

Line 63, replace "NPVAT" with --NPVAT--;

Line 64, replace "LQRCN" with --LQRCN--;

Line 65, replace "NVDFN" with --NVDFN--.

Column 10, Line 19, replace "2.2 kb" with --2.2 kb EcoRI-BspMI
fragment from pUC-Av2 and inserting it into the EcoRI cleavage
site of pUC19. The PCR products are then amplified in bacteria
and sequenced, and the 1.4 kb EcoNI-XcmI fragment which contains
P1 is subcloned in pUC-AV2 in which the corresponding wild-type
cap sequence has been cut out. Accordingly, the plasmids

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(mutants) called after the AA insertion sites pI-447, PI-534, pI-573 and pI-587 contained the complete AAV2 genome.--

Column 17, Claim 2, Line 23, replace "insertions" with
--insertion--.

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